Raw Sequence Listing Error Summary

	ERROR DETECTED	SUGGESTED CORRECTION	SERIAL NUMBER:	SERIAL NUMBER:	
		EASE DISREGARD ENGLISH "ALPHA" HEADERS, V	VHICH WERE INSERTED BY PTO S	OFTWARE	
ATTN:		The number/text at the end of each line "wrapped" down	to the next line.		
1	Wrapped Nucleics	This may occur if your file was retrieved in a word proce-	ssor after creating it.	RECEIVED	
		Please adjust your right margin to .3, as this will preven	"wrapping".	H L O L 10-22	
		Please adjust your right margin to .o, as the will prove		00	
		The amino acid number/text at the end of each line "wra	noed " down to the next line.	SEP 0.6 2000	
2	Wrapped Aminos	This may occur if your file was retrieved in a word proce	essor after creating it.	JEI OO	
		This may occur if your file was retrieved in a word prove	t "wranning"	-44	
		Please adjust your right margin to .3, as this will preven	. wapping .	TECH CENTER 1000/2900	
		#	longth. This includes snaces	(EOH OFILE)	
3	Incorrect Line Length	The rules require that a line not exceed 72 characters in	length. This includes spaces.		
	(·		This may be caused by the use of to	abs	
4	Misaligned Amino Acid	The numbering under each 5th amino acid is misaligned	tehe and use spacing between the fi	ímbers -	
	Numbering	between the numbering. It is recommended to delete an	y tabs and use spacing between the, in		
			U. O Dulas		
5	Non-ASCII	This file was not saved in ASCII (DOS) text, as required	by the Sequence Rules.	1	
		Please ensure your subsequent submission is saved in	ASCII text so that it can be processed	1.	
6	Variable Length	Sequence(s) contain n's or Xaa's which represent	ed more than one residue.		
`		As per the rules, each n or Xaa can only represent a sin	igle residue.		
		Please present the maximum number of each residue h	aving variable length and		
		indicate in the (ix) feature section that some may be mis	ssing.		
-	Detections 2.0 "bug"	A "bug" in Patentin version 2.0 has caused the <220>-	<223> section to be missing from amir	no acid	
′—	Patentin ver. 2.0 "bug"	sequence(s)	automatically generate this section from	om the	
		previously coded nucleic acid sequence. Please manu	ally copy the relevant <220>-<223> se	ection	
		to the subsequent amino acid sequence. This applies	primarily to the mandatory <220>	-<223>	
		to the subsequent animo dot sequences. The spring		-4-	
		sections for Artificial or Unknown sequences.			
		Sequence(s) missing. If intentional, please use the	e following format for each skipped se	equence:	
8	Skipped Sequences	Sequence(s)missing. If intentional, please use to	e following formactor outsit supplies	•	
	(OLD RULES)	(2) INFORMATION FOR SEQ ID NO:X: (i) SEQUENCE CHARACTERISTICS:(Do not insert a	ny headings under "SEQUENCE CH/	ARACTERISTICS")	
		(i) SEQUENCE CHARACTERISTICS:(Uo not insert a	ny headings under Seasched St.	"""""""""""""""""""""""""""""""""""""""	
		(xi) SEQUENCE DESCRIPTION:SEQ ID NO:X:			
		This sequence is intentionally skipped			
			an a latata tha aldonod o	naunno(s)	
		Please also adjust the "(iii) NUMBER OF SEQUENCE	S:" response to include the skipped so	equence(s).	
			s u i - ft f-r each akinned se	nguence	
9	Skipped Sequences	Sequence(s) missing. If intentional, please use the	e following format for each shipped se	Addenoc.	
	(NEW RULES)	<210> sequence id number			
	,	<400> sequence id number			
		000			
10	Use of n's or Xaa's	Use of n's and/or Xaa's have been detected in the Seq	uence Listing.		
	(NEW RULES)	Use of <220> to <223> is MANDATORY if n's or Xaa's	s are present.	4.	
	(11211110220)	In <220> to <223> section, please explain location of n	or Xaa, and which residue n or Xaa	represents.	
4.4	Use of <213>Organism	Sequence(s) are missing this mandatory field	or its response.		
''	/ (NEW RULES)	004001100(0)		-	
,	(NEW ROLES)	·			
/	Harard 2000s Facilities	Sequence(s) are missing the <220>Feature and	associated headings.		
12	Use of <220>Feature	Use of <220> to <223> is MANDATORY if <213>OR	GANISM is."Ártificial" or "Unknown"	•	
	(NEW RULES)	Please explain source of genetic material in <220>	to <223> section.		
		(See "Federal Register," 6/01/98, Vol. 63,	No. 104 pp. 29631-32) (Sc	ec 1 823 of new Rules)	
		(See "Federal Register," 6/01/90, Vol. 63,	140. 104, pp. 20001 02) (00		
				runted	
13	Patentin ver. 2.0 "bug"	Please do not use "Copy to Disk" function of Pate	entin version 2.0. This causes a corr	upieu	
	_	file, resulting in missing mandatory numeric identifiers	and responses (as indicated on raw s	equence listing).	
		Instead, please use "File Manager" or any other means	s to copy file to floppy disk.	•	

Application No. 09/544045

NOTICE TO COMPLY WITH REQUIREMENTS FOR PATENT APPLICATIONS CONTAINING NUCLEOTIDE SEQUENCE AND/OR AMINO ACID SEQUENCE DISCLOSURES

The nucleotide and/or amino acid sequence disclosure contained in this application does

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not comply with the requirements for such a disclosure as set forth in 37 CFR 1.821 -1.825 for the following reason(s): This application clearly fails to comply with the requirements of 37 CFR 1.821 - 1.825. Applicant's attention is directed to these regulations, published at 1114 OG 29, May 15, 1990 and at 55 FR 18230, May 1, 1990. 2. This application does not contain, as a separate part of the disclosure on paper copy, a "Sequence Listing" as required by 37 CFR 1.821(c). A copy of the "Sequence Listing" in computer readable form has not been submitted as required by 37 CFR 1.821(e). A copy of the "Sequence Listing" in computer readable form has been submitted. However, the content of the computer readable form does not comply with the requirements of 37 CFR 1.822 and/or 1.823, as indicated on the attached copy of the marked-up "Raw Sequence Listing." The computer readable form that has been filed with this application has been found to be damaged and/or unreadable as indicated on the attached CRF Diskette Problem Report. A substitute computer readable form must be submitted as required by 37 CFR 1.825(d). The paper copy of the "Sequence Listing" is not the same as the computer readable form of the "Sequence Listing" as required by 37 CFR 1.821(e). 7. Applicant must provide: An initial or substitute computer readable form (CRF) copy of the "Sequence Listing" An initial or substitute paper copy of the "Sequence Listing", as well as an amendment directing its entry into the specification A statement that the content of the paper and computer readable copies are the same and, where applicable, include no new matter, as required by 37 CFR 1.821(e) or 1.821(f) or 1.821(g) or 1.825(b) or 1.825(d)

Please return a copy of this notice with your response.

For Rules Interpretation, call (703) 308-1123 For CRF submission help, call (703) 308-4212 For PatentIn software help, call (703) 557-0400

For questions regarding compliance with these requirements, please contact: